都道府県薬剤師会担当役員殿

日本薬剤師会 担当副会長 川上 純一

第十八改正日本薬局方(英文版)正誤表の送付について(その3)

平素より、本会会務に格別のご高配を賜り厚く御礼申し上げます。

標記について、厚生労働省医薬局医薬品審査管理課より、別添のとおり事務連絡がありましたのでお知らせいたします。

会務ご多用のところ恐縮ながら、貴会会員にご周知下さるようお願い申し上げます。



事 務 連 絡 令和5年11月10日

公益社団法人日本薬剤師会 御中

厚生労働省医薬局医薬品審査管理課

第十八改正日本薬局方(英文版)正誤表の送付について(その3)

標記について、別添写しのとおり各都道府県衛生主管部(局)薬務主管課宛に連絡しましたので、お知らせいたします。



事 務 連 絡 令和5年11月10日

各都道府県衛生主管部(局)薬務主管課 御中

厚生労働省医薬局医薬品審査管理課

第十八改正日本薬局方(英文版)正誤表の送付について「(その3)

第十八改正日本薬局方(令和3年厚生労働省告示第220号)の英文版につきまして、 一部に誤植等がありましたので別紙のとおり正誤表を送付いたします。 General Tests / 1.09 Qualitative Tests

residue in diluted and filter if

General Tests / 7.03 Test for Rubber Closure for Aqueous Infusions

Page	Line	Correction	Error
	left ↓17	Further, to exactly 1 mL of Standard Zinc	Further, to exactly 1 mL of Standard Zinc
		Solution for atomic absorption	Solution for atomic absorption
202		spectrophotometry add diluted dilute nitric	spectrophotometry add diluted nitric acid (1 in
		acid (1 in 3) to make exactly 20 mL, and use	3) to make exactly 20 mL, and use this
,		this solution as the standard solution.	solution as the standard solution.

General Tests / 9.22 Standard Solutions

Page	Line	Correction	Епог
219	left ↑ 21-23	Standard Cadmium Solution Measure exactly 10 mL of Standard Cadmium Stock Solution, and add diluted dilute nitric acid (1 in 3) to make exactly 1000 mL. Pipet 10 mL of this solution, and add diluted dilute nitric acid (1 in 3) to make 100 mL. Each mL of this solution contains 0.001 mg of cadmium (Cd). Prepare before use.	Standard Cadmium Solution Measure exactly 10 mL of Standard Cadmium Stock Solution, and add diluted nitric acid (1 in 3) to make exactly 1000 mL. Pipet 10 mL of this solution, and add diluted nitric acid (1 in 3) to make 100 mL. Each mL of this solution contains 0.001 mg of cadmium (Cd). Prepare before use.

Official Monographs

Aminophylline Hydrate アミノフィリン水和物

Page	Line	Correction	Error
448	right ↓5	(C7H8N4O2)2 · C2H8N2 · xH2O	<u>C14H16N8O4.</u> C2H8N2. xH2O

L-Aspartic Acid L-アスパラギン酸

Page	Line	Correction	Епог
		(3) Sulfate <1.14>—Dissolve 0.6 g of	(3) Sulfate <1.14>—Dissolve 0.6 g of
		L-Aspartic Acid in 5 mL of dilute hydrochloric	L-Aspartic Acid in 5 mL of dilute hydrochloric
	right ↑19	acid and 30 mL of water, add water to make 45	acid and 30 mL of water, add water to make 45
487		mL, and add 5 mL of barium chloride TS.	mL, and add 5 mL of barium chloride TS.
		Perform the test with this solution as the test	Perform the test with this solution as the test
		solution. Prepare the control solution with 0.35	solution. Prepare the control solution with 0.35
		mL of 0.005 mol/L sulfuric acid VS, add 5 mL	mL of 0.005 mol/L sulfuric acid VS, add 5 mL
		of dilute hydrochloric acid and water to make	of dilute hydrochloric acid and water to make
		45 mL, and add 5 mL of barium chloride TS	45 mL, and add 5 mL of barium chloride (not
		(not more than 0.028%).	more than 0.028%).

Bicalutamide ビカルタミド

icalutamide E	コルタミト		
Page	Line	Correction	Error
		For the areas of the peaks, related substance G,	For the areas of the peaks, related substance G,
		having the relative retention times of about	having the relative retention times of about
		0.21 and about 0.25, related substance I,	0.21 and about 0.25, related substance I,
		having the relative retention time of about	having the relative retention time of about
		0.23, related substance M, related substance N,	0.23, related substance M, related substance N,
		related substance O, having the relative	related substance O, having the relative
550	left ↑4	retention time of about 0.55, related substance	retention time of about 0.55, related substance
		A, having the relative retention time of about	A, having the relative retention time of about
-		0.95, and related substance K, and related	0.95, and related substance L, and related
v		substance P, having the relative retention time	substance P, having the relative retention time
		of about 1.09 from the sample solution,	of about 1.09 from the sample solution,
		multiply their correction factors, 0.5, 0.5, 0.5,	multiply their correction factors, 0.5, 0.5, 0.5,
		0.4, 0.7, 0.5, 1.1, 0.9 and 0.7, respectively.	0.4, 0.7, 0.5, 1.1, 0.9 and 0.7, respectively.

Ciprofloxacin Hydrochloride Hydrate シプロフロキサシン塩酸塩水和物

Cipionoxaciii 11	diocinoride 11)	Tate - Introduction of the party	
Page	Line	Correction	Error
765	left ↓8	[86393-32-0, monohydrate]	[86393-32-0, monohydrochloride monohydrate]

Clotrimazole クロトリマゾール

Page	Line	Correction	Еггог
		(3) Sulfate <1.14>—Dissolve 0.5 g of	(3) Sulfate <1.14>—Dissolve 0.5 g of
		Clotrimazole in 10 mL of methanol, and add 1	Clotrimazole in 10 mL of methanol, and add 1
		mL of dilute hydrochloric acid and water to	mL of dilute hydrochloric acid and water to
799		make 50 mL. Perform the test using this	make 50 mL. Perform the test using this
	right ↑9	solution as the test solution. Prepare the	solution as the test solution. Prepare the
		control solution with <u>0.50</u> mL of 0.005 mol/L	control solution with <u>0.05</u> mL of 0.005 mol/L
		sulfuric acid VS, 10 mL of methanol, 1 mL of	sulfuric acid VS, 10 mL of methanol, 1 mL of
		dilute hydrochloric acid and water to make 50	dilute hydrochloric acid and water to make 50
		mL (not more than 0.048%).	mL (not more than 0.048%).

Fursultiamine Hydrochloride フルスルチアミン塩酸塩

Page	Line	Correction	Error
1051	right ↓ 27	[2105-43-3]	[804-30-8, Fursultiamine]

Glycerin グリセリン

·	IJ COLIN	C / V		a contract of the contract of
	Page	Line	Correction	Еггог
	1000	1.0 14	Description Glycerin is a clear, colorless,	Description Glycerin is a clear, colorless,
	1080	left ↓14	viscous liquid.	viscous liquid. It has a sweet taste.

Dental Iodine Glycerin 歯科用ヨード・グリセリン

ental fodine Glycerin 国行力コート・クリビリン				
Page	Line	Correction	Ептог	
1173	left ↓24	(2) Potassium iodide—Separate the water layers of the sample solution and standard solution obtained in (1), pipet 7mL each of the water layers, and to each add exactly 1mL of diluted dilute hydrochloric acid (1 in 2), 1 mL of sodium nitrite TS and 10 mL of a mixture of chloroform and hexane (2:1), and shake immediately.	(2) Potassium iodide—Separate the water layers of the sample solution and standard solution obtained in (1), pipet 7mL each of the water layers, and to each add exactly 1mL of diluted hydrochloric acid (1 in 2), 1 mL of sodium nitrite TS and 10 mL of a mixture of chloroform and hexane (2:1), and shake immediately.	

Ketoprofen ケトプロフェン

stephoten / 1/2-2-0				
Page	Line	Correction	Епог	
1224		Control solution: To a mixture of 0.6 mL of	Control solution: To a mixure of 0.6 mL of	
		Cobalt (II) Chloride CS and 2.4 mL of Iron	Cobalt (II) Chloride CS and 2.4 mL of Iron	
	right ↑	(III) Chloride CS add diluted dilute	(III) Chloride CS add diluted hydrochloric acid	
	20,21,23	hydrochloric acid (1 in 10) to make 10 mL. To	(1 in 10) to make 10 mL. To 5.0 mL of this	
		5.0 mL of this solution add diluted dilute	solution add diluted hydrochloric acid (1 in 10)	
		hydrochloric acid (1 in 10) to make 100 mL.	to make 100 mL.	

Loxoprofen Sodium Hydrate ロキソプロフェンナトリウム水和物

Page	Line	Correction	Error
1279	right ↓ 17	[<u>226721-96-6</u>]	[<u>80382-23-6</u>]

Miconazole ミコナゾール

Page	Line	Correction	Error
1255		Loss on drying <2.41> Not more than 0.5% (1	Loss on drying <2.41> Not more than 0.5% (1
1357		g, in vacuum, silica gel, 60°C, 3 hours).	g, in vacuum, silica gel, 60%, 3 hours).

Mosapride Citrate Tablets モサプリドクエン酸塩錠

Page	Line	Correction .	Етгот
	ľ	Add 9 mL of methanol, shake for 20 minutes,	Add 9 mL of methanol, shake for 20 minutes,
20		centrifuge, and use the supernatant liquid as	centrifuge, and use the supernatant liquid as
		the sample solution. Pipet 1 mL of this	the sample solution. Pipet 1 mL of this
1389	right ↓5	solution, add methanol to make exactly 20 mL.	solution, add methanol to make exactly 20 mL.
		Pipet 2 mL of this solution, add methanol to	Pipet 2 mL of the sample solution, add
		make exactly 20 mL, and use this solution as	methanol to make exactly 20 mL, and use this
		the standard solution.	solution as the standard solution.

Pitavastatin Calcium Hydrate ピタバスタチンカルシウム水和物

Page	Line	Correction	Error
		The control solution is prepared as follows:	The control solution is prepared as follows:
		Take 10 mL of a solution of magnesium nitrate	Take 10 mL of a solution of magnesium nitrate
		hexahydrate in ethanol (95) (1 in 10), and fire	hexahydrate in ethanol (95) (1 in 10), and fire
1540	right ↓5	the ethanol to burn. Hereafter, proceed as for	the ethanol to burn. Hereafter, proceed as for
		the test solution, then add 2.0 mL of Standard	the test solution, then add 2.0 mL of Standard
		Lead Solution, 2 mL of dilute acetic acid and	Lead Solution, 2 mL of acetic acid and water
		water to make 50 mL (not more than 20 ppm).	to make 50 mL (not more than 20 ppm).

Pitavastatin Calcium Tablets ピタバスタチンカルシウム錠

Page	Line	Correction	Error
21.2		6-{2-[2-Cyclopropyl-4-(4-fluorophenyl)quinol	6-{2-[2-cyclopropyl-4-(4-fluorophenyl)quinoli
1545	left ↓ 1-2	in-	n-
		3-yl]ethenyl}-4-hydroxyoxane-2-one	3-yl]ethenyl}-4-hydroxyoxane-2-one

D-Sorbitol D-ソルビトール

Page	Line	Correction	Ептог
		(7) Glucose—Dissolve 20.0 g of D-Sorbitol in	(7) Glucose—Dissolve 20.0 g of D-Sorbitol in
		25 mL of water, and boil gently with 40 mL of	25 mL of water, and boil gently with 40 mL of
		Fehling's TS for 3 minutes. After cooling, filter	Fehling's TS for 3 minutes. After cooling, filter
	right ↓ 10-11	the supernatant liquid cautiously through a	the supernatant liquid cautiously through a
1733		glass filter (G4), leaving the precipitate in the	glass filter (G4), leaving the precipitate in the
F 0 ga		flask as much as possible, wash the precipitate	flask as much as possible, wash the precipitate
		with hot water until the last washings no	with hot water until the last washings no
		longer show alkalinity, and filter the washings	longer show an alkali reaction, and filter the
		through the glass filter.	washings through the glass filter.

Voglibose ボグリボース

Page	Line	Correction	Error
1911	left ↑25	It is very soluble in water, freely soluble in acetic acid (100), slightly soluble in methanol, and very slightly soluble in ethanol (99.5).	It is very slightly soluble in water, freely soluble in acetic acid (100), slightly soluble in methanol, and very slightly soluble in ethanol (99.5).

Zopiclone ゾピクロン

Page	Line	Correction	Error
55.7	right ↓ 33-36	determine each peak area by the automatic integration method: the peak areas of related substance A, having the relative retention time of about 0.1 to zopiclone, related substance B, having the relative retention time of about 0.2, related substance C, having the relative retention time of about 0.5, related substance D, having the relative retention time of about 0.9, obtained from the sample solution are not larger than 1/10 times the peak area of zopiclone from the standard solution, and the peaks mentioned above from the sample solution is not larger than 1/10 times the peak area of zopiclone from the standard solution.	determine each peak area by the automatic integration method: the peak areas of related substance A, having the relative retention time of about 0.1 to zopiclone, related substance B, having the relative retention time of about 0.2, related substance C, having the relative retention time of about 0.5, related substance D, having the relative retention time of about 0.9 and the peaks other than mentioned above, obtained from the sample solution, are not larger than 1/10 times the peak area of zopiclone from the standard solution.